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Non-Maxwellian viscoelastic stress relaxations in soft matter†

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Viscoelastic stress relaxation is a basic characteristic of soft matter systems such as colloids, gels, and biological networks. Although the Maxwell model of linear viscoelasticity provides a classical description of stress relaxation, it is often not sufficient for capturing the complex relaxation dynamics of soft matter. In this Tutorial, we introduce and discuss the physics of non-Maxwellian linear stress relaxation as observed in soft materials, the ascribed origins of this effect in different systems, and appropriate models that can be used to capture this relaxation behavior. We provide a basic toolkit that can assist the understanding and modeling of the mechanical relaxation of soft materials for diverse applications.

Soft matter systems are characterized by dynamic fluctuations and rearrangements within the microstructure, which play an important role in the function of a wide range of soft materials, such as cell-matrix interactions in biology,^{1,2} or energy dissipation and self-healing in engineered soft materials.^{3,4} These rearrangement events give rise to a time-dependent response in the mechanical properties, *i.e.*, viscoelasticity.⁵ The measurement

of viscoelasticity can thus provide meaningful information into the dynamics of the soft material of interest. This is most commonly done at bulk scales using a dynamic mechanical analyzer⁶ or a rheometer,⁵ though microscopic-scale measurements can also be made *via* microrheology⁷ and scattering.⁸

The dynamics of soft matter systems are non-trivial, and are characterized by a viscoelastic response that is often more complex than the Maxwell model of linear viscoelasticity (hence the term “complex fluids”). The Maxwell model is a canonical model introduced in elementary studies of linear viscoelasticity, which is characterized by a single characteristic relaxation time. However, measurements on real soft matter systems commonly exhibit a viscoelastic response that underscores a broad distribution of relaxation modes. These distributions of relaxation processes are generally interpreted as arising from a wide range of structural length-scales or relaxation mechanisms by which stress can relax, though the exact origins of these mechanisms are often not clear. Indeed, there is a vast

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where η is the (linear) Newtonian viscosity of the dashpot, G is the (linear) Hookean elasticity of the spring, and their ratio describes a characteristic relaxation time $\eta/G = \tau_c$. The Maxwell viscoelastic model is often used to model stress relaxation processes arising from step strain perturbations, $\gamma(t) = \gamma_0 \mathcal{H}(t)$ (where $\mathcal{H}(t)$ is the Heaviside step function), or small-amplitude oscillatory strain perturbations, $\gamma(t) = \gamma_0 \sin(\omega t)$ (Fig. 1B and C). Solving the constitutive relation in the case of a step strain experiment, we find that the relaxation modulus $G(t)$ of a soft material system described by eqn (1) exhibits an exponential decay (Fig. 1D):

$$G(t) = G_0 \exp(-t/\tau_c) \quad (2)$$

where G_0 is the plateau modulus of the material. Solving the constitutive relation for the Maxwell model in the case of a linear oscillatory strain, the storage $G'(\omega)$ and loss $G''(\omega)$ components of the complex shear modulus $G^*(\omega)$ have the forms (Fig. 1E):

$$G^*(\omega) = G'(\omega) + \mathrm{i}G''(\omega) = \frac{G_0}{1 + \mathrm{i}\omega\tau_c} \quad (3)$$

or by separating the real and imaginary components:

$$\begin{aligned} G'(\omega) &= G_0 \frac{(\omega \tau_c)^2}{1 + (\omega \tau_c)^2} \\ G''(\omega) &= G_0 \frac{\omega \tau_c}{1 + (\omega \tau_c)^2} \end{aligned} \quad (4)$$

These materials functions exhibit a characteristic cross-over at a frequency of $\omega_c = 1/\tau_c$, with terminal power-law slopes at low frequency ($\omega \ll \omega_c$) of 2 and 1 for the storage and loss moduli, respectively. The Maxwell model response for oscillatory strain is thus dictated by the Deborah number arising from the oscillatory strain protocol, $De = \omega\tau_c$. The Deborah number helps characterize the importance of linear viscoelastic effects in the system at different deformation rates, with $De \ll 1$ indicating a viscous liquid-dominated response, $De \gg 1$ indicating an elastic solid-dominated response, and $De = 1$ indicating the point of cross-over between the two response regimes.

In addition to the typical frequency sweep plot showing the storage and loss moduli as a function of frequency (Fig. 1E), there are other methods for representing $G'(\omega)$ and $G''(\omega)$ data

I. Maxwell model of linear viscoelasticity

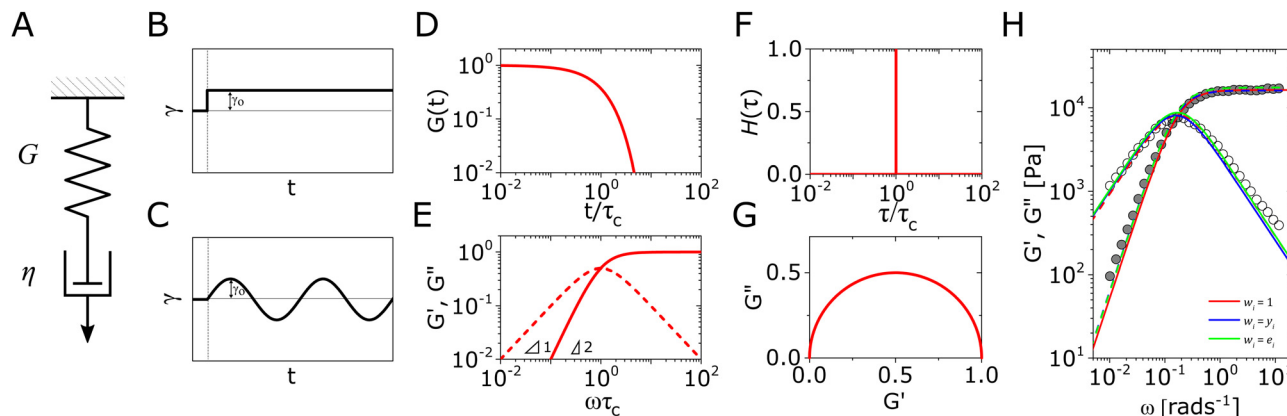
Exponential decays of the form $\phi = \phi_0 \exp(-t/\tau_c)$, where ϕ is an observable thermo-physical property and ϕ_0 is the initial value at $t = 0$, are often used to model relaxation events arising from simple dynamical processes. For instance, exponential decays are exact solutions to describe the correlation of Brownian motion of particles, the correlation of dipoles in the rotational



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Yet another useful representation for describing the viscoelasticity of soft matter is the continuous relaxation spectrum $H(\tau)$, which encodes the distribution of relaxation modes in a given system (more detail in Section III).^{15,16} The continuous relaxation spectrum is encoded in the hereditary integral formulation of linear viscoelasticity (*via* the Boltzmann superposition principle),¹⁷ and describes the relaxation modulus *via*:

$$G(t) = \int_{-\infty}^{\infty} H(\tau) \exp(-t/\tau) d \ln \tau \quad (5)$$

$$\begin{aligned} G'(\omega) &= \int_{-\infty}^{\infty} H(\tau) \frac{(\omega\tau)^2}{1 + (\omega\tau)^2} d\ln \tau \\ G''(\omega) &= \int_{-\infty}^{\infty} H(\tau) \frac{\omega\tau}{1 + (\omega\tau)^2} d\ln \tau \end{aligned} \quad (6)$$

Finally, in response to a step stress of amplitude σ_0 , the Maxwell model exhibits an instantaneous increase in creep compliance $J(t) = \gamma(t)/\sigma_0$ (thus exhibiting a retardation time of zero), followed by a linear increase in creep compliance $J(t)$ with time that is given by the relation:

$$J(t) = \frac{1}{G} + \frac{t}{\eta} \quad (7)$$

Though most viscoelastic stress relaxation responses of soft materials are more complex than a single exponential decay (eqn (2)), there are a few exceptional cases where essentially Maxwellian behavior is observed. A prototypical example of materials exhibiting Maxwellian viscoelasticity are transient polymer networks, such as those consisting of star poly(ethylene glycol) (PEG) which are end-functionalized with diverse binding

A. Non-first-order reaction kinetics

Deviations from Maxwellian viscoelastic relaxation can arise in telechelic networks when the bond interactions between polymers do not follow first-order reaction kinetics. A representative example of these kinds of interactions come from host-guest interactions, which are often structurally complex and can give rise to a complex dissociation pathway with many intermediate steps.⁴⁸ Indeed, studies have shown that multi-arm PEGs functionalized with cyclodextrin and cholesterol exhibit strong deviations from Maxwellian rheology,⁴⁹ in contrast to multi-arm PEGs functionalized with dynamic covalent linkages which exhibit Maxwellian linear viscoelastic responses.^{21,25,50} The use of such complex interaction moieties provides an interesting, chemically-oriented approach to tuning the relaxation spectrum of the polymeric network.

B. Chain dynamics

Deviations from single-mode Maxwell viscoelastic responses are also expected based on classical models of polymer dynamics. In the Rouse model, the relaxation dynamics are assumed to arise from the Brownian motion of beads connected by springs, wherein the terminal relaxation time depends on the polymer chain length. This leads to a stress relaxation response arising from a linear summation of individual (exponential) Rouse relaxation modes, which results in a power-law scaling of the relaxation modulus preceding a terminal relaxation, of the form:⁵¹

$$G(t) \sim G_0(t/\tau_0)^{-1/2} \exp(-t/\tau_R) \quad (8)$$

where τ_0 is the shortest relaxation mode (of an individual Rouse segment) and τ_R is the longest relaxation mode of the entire chain. The Zimm model includes additional hydrodynamic interactions between individual segments of the bead-spring chain (and thus is more appropriate for dilute solutions of macromolecules where hydrodynamic effects are not screened). This changes the power-law scaling in eqn (8) from $\sim t^{-1/2}$ to $\sim t^{-1/3\nu}$ where ν is the Flory scaling exponent that is related to the solvent quality.⁵¹

Non-Maxwellian stress relaxation becomes more prominent for polymer melts with long entangled chains, in which Rouse and Zimm modes are followed at longer times by entanglement effects.⁵² This can result in the addition of stress relaxation mechanisms, such as reptation, contour-length fluctuations and constraint release.⁵³ More complex relaxation processes can also arise specifically due to entanglement effects in polymer systems of more complex topologies, such as entangled star polymers⁵⁴ and entangled ring polymers.⁵⁵

Semiflexible polymers such as cytoskeletal polymers and extracellular matrix polymers have been shown to exhibit unique chain relaxation dynamics – distinct from Rouse and Zimm dynamics of polymers – which arise due to the significant thermal undulations of the semiflexible polymer backbone. Studies have shown that in such systems, the viscoelastic moduli exhibit a power-law scaling of $G'(\omega) \sim G''(\omega) \sim \omega^n$, in which $n = 3/4$ in the high frequency regime.^{56–58}

C. Sticky chain dynamics

Though Rouse dynamics are expected to occur even in swollen gels such as the Maxwellian transient networks introduced in Section I, these relaxation modes occur at much shorter time-scales than those arising from the reversible interactions in these systems, and are often not measurable. However, when there is a sufficient concentration of interacting stickers per polymer chain in such transient networks, we may instead observe the emergence of sticky chain dynamics.

One such manifestation is the sticky Rouse model,^{36,40,59} in which the associations between neighboring polymer chains are understood to constrain the polymer chains and lead to a Rouse-like dynamical regime at long times. This results in the observation of power-law relaxation of the form $G(t) \sim t^{-1/2}$ in associative polymers, most commonly in systems where the main chain has been functionalized by binding motifs such that multiple stickers exist per chain.^{59–62} In similar vein, the concept of sticky dynamics have also been applied to reptation dynamics for studying associative networks with longer entangled chains.^{63–65}

Sticky dynamics can also give rise to non-Maxwellian viscoelasticity in semiflexible polymer networks. Modeling studies have shown that dissociation events along the backbone of a semiflexible polymer generate transverse relaxation modes, in which successive relaxation modes become progressively slower as more dissociation events need to occur; this leads to a terminal power-law relaxation of $G(t) \sim t^{-1/2}$, which is commonly observed in semiflexible polymer networks.⁶⁶ Experimentally, this power-law region has also been shown to be characterized by stress fluctuations indicative of collective dynamics.⁶⁷

D. Multiple relaxation modes

Non-Maxwellian viscoelastic responses are a natural outcome for systems which intrinsically have multiple relaxation modes, such as those with chain length polydispersity, sticker heterogeneity, or multiple relaxation mechanisms.

Polydispersity in the chain length can result in heterogeneous chain dynamics, and thus a broad distribution of relaxation modes. The observation of non-Maxwellian stress relaxations can be leveraged to gain insight into the polydispersity of polymer systems. For instance, for melt systems undergoing reptation, the observed distribution of relaxation modes (from $H(\tau)$) can be retraced to the molecular weight distribution of the system.⁶⁸

For associative polymers, multiple relaxation modes can arise due to factors such as the sticker distribution on chains⁶⁹ and sticker clustering.⁷⁰ Stickers can also introduce spatial mismatches in the binding of two chains, resulting in formation of defects such as chain loops. This can generate energetic penalties in the chain and cause broadening in the viscoelastic relaxation curve.⁷¹ These effects – especially in tandem with chain length polydispersity – can lead to complicated linear viscoelastic responses which can be challenging to ascribe to a single factor.

For composite materials such as polymer-particle systems, non-Maxwellian viscoelastic responses can arise due to multiple relaxation mechanisms that arise intrinsically from the composite microstructure. For instance, latex particles which



are bridged by HEUR micelles exhibit a composite response of Rouse dynamics, bridging interactions, and large-scale cluster dynamics, the sum of which can lead to complex non-Maxwellian viscoelastic behavior.^{72,73} Furthermore, each of these relaxation mechanisms can be governed by a distribution of relaxation modes. For the contribution arising from Rouse dynamics, such distribution can arise as a result of chain polydispersity. For the bridging dynamics, it is well understood that the number of bridging linkers can significantly change the terminal relaxation time of a pair of particles due to cooperativity.⁷⁴ As such, a Poisson distribution of linkers in associating particle systems can lead to a significantly non-Maxwellian viscoelastic response, which can be useful for understanding the dynamics of particle-polymer systems such as latex and HEUR, metal-coordinating nanoparticle hydrogels and coordination cage hydrogels.⁷⁵ Finally, cluster dynamics can also be influenced by a distribution in cluster sizes, which we treat separately in Section III E.

Lastly, a broad distribution in relaxation modes can arise in systems which are characterized by a distribution of activation energies. For instance, a Gaussian distribution in the activation energy of relaxation can facilitate a log-normal distribution in the stress relaxation time (see Section IV A). This approach has been taken to model the relaxation of a metal-coordinate polymer material⁷⁶ (in which bond strengths are assumed to be distributed in a Gaussian manner) as well as glassy systems (in which relaxation is assumed to arise from local domains with a Gaussian distribution of activation energies).^{77,78}

E. Convolution of relaxation processes

A stretched exponential relaxation of the form

$$G(t) = G_0 \exp[-(t/\tau_c)^\beta] \quad (9)$$

is observed ubiquitously in soft materials such as glasses,^{79–81} gels,⁸² tissues,⁸³ and surfactants.²⁹ Several studies have shown that this stretched exponential relaxation function is a direct outcome of relaxation processes which are convolved by a structural distribution of the relaxing unit. Mathematically, this can be expressed by an integral convolution of relaxation processes, such that:⁸⁴

$$G(t) = G_0 \int_0^\infty \Phi(\tau) Q(t, \tau) d\tau \quad (10)$$

where $Q(t, \tau)$ is a function describing the relaxation, and $\Phi(\tau)$ is a probability function for a given relaxation time τ that is convolved with $Q(t, \tau)$. In the framework of soft matter relaxation, $Q(t, \tau)$ directly describes the exponential relaxation of the primary unit of a given system (e.g., reptation time of a chain or relaxation time of a cluster), and $\Phi(\tau)$ is related to a distribution in the topological features such as the length or size of the relaxing unit, which weights the size-dependent relaxation function $Q(t, \tau)$.

An example of this idea is the work of Douglas *et al.*,^{85–87} which evaluates the relaxation function of polymeric systems that form clusters. In these works, it is assumed that the cluster relaxation time depends on diffusion processes that scale with the size of the cluster. It is thus shown that the relaxation time

scales as $\tau \sim L^2/D_0$ with L being the characteristic length of the cluster and D_0 being the diffusivity of the cluster. Assuming that clusters take a Boltzmann (exponential) size distribution, one can derive the convolution function $\Phi(\tau) = \exp(-L) = \exp(-\tau^{1/2})$. A steepest descent approximation of eqn (10) with this convolution function results in the following derivation:

$$G(t) = G_0 \int_0^\infty \exp(-\tau^A) \exp(t/\tau) d\tau = G_0 \exp(-(t/\tau)^\beta) \quad (11)$$

where $A = 1/2$ and the stretching exponent $\beta = A/(A + 1) = 1/3$.⁸⁴ Thus, cluster size polydispersity itself can lead to a stretched exponential of the form $G(t) \sim \exp(-(t/\tau)^{1/3})$, a form commonly seen in the stress relaxation of polymeric systems such as gels.^{85–87}

A similar derivation has been proposed by Cates *et al.* for wormlike micelles undergoing reptation,²⁹ with the main difference being an additional assumption that the diffusivity $D_0 \sim 1/L$ arises due to the curvilinear diffusion of the chains along their confining tubes. The convolution function thus becomes $\Phi(\tau) = \exp(-L) = \exp(-\tau^{1/3})$, resulting in a final stretched exponential function of $G(t) \sim \exp(-(t/\tau)^{1/4})$. Overall, the examples by Douglas *et al.*, and Cates *et al.*, show that the stretched exponential stress relaxation is a mathematical outcome of convoluted relaxation events in which there is a size polydispersity of the relaxing units, for instance clusters or tubes.

Finally, a related approach is the work of Curro *et al.*,⁸⁸ who have shown that permanently cross-linked systems such as elastomers can also exhibit a power-law stress relaxation. This relaxation is ascribed to dangling chain ends in the system, such that $G(t) \sim \sum_{k=1}^\infty \sigma_k(t) P(k)$, where k is the length of the chain

branch, $\sigma_k(t) = k - l(t)$ describes the stress arising from parts of the dangling chains which have not relaxed ($l(t)$ is the lognormally-dependent relaxation rate of dangling chain ends as derived by de Gennes), and $P(k)$ is the probability of having a dangling chain of length k . The convolution of the probability of k by the time-dependent stress arising from k is shown to result in a power law relaxation $G(t) \sim t^{-(q/a)}$ where q is the ratio of cross-link density to monomer density, and a is a material constant related to the reptation time of the chain.⁸⁸ This idea of the disentanglement of dangling chain ends has also been utilized by Rubinstein *et al.* to explain the power-law stress relaxation in block copolymers, see ref. 89.

F. Criticality and fractals

The rheological response of an associating soft material near critical points are often characterized by a power-law response in the frequency-dependent viscoelastic moduli such that $G'(\omega) \sim G''(\omega) \sim \omega^n$. This is observed in cross-linking polymers approaching gelation,⁹⁰ in soft colloidal systems approaching jamming,⁹¹ and in fiber networks approaching critical connectivities.⁹² Though these studies have canonically predicted $n = 1/2$ across the different systems, the actual exponent underlying the power-law relaxation has been shown to vary significantly depending on the specific nature of the system.



One important factor that appears to govern n is the underlying fractal structure of the associated material. In branched polymeric materials, Muthukumar *et al.* has shown that n can be related to the fractal dimension d_f of the polymer network that forms at the percolation threshold.^{93–95} Other works have shown that the fractal nature of the percolating system remains in the gel structure well beyond the percolation threshold, and that these structures result in remnant signatures in the loss modulus⁹⁶ and the relaxation spectrum of the resulting gel.^{97,98} In marginal spring networks, the underlying fractal structure has also been associated with a value of n that is substantially lower than $1/2$ at high frequencies,⁹⁹ though a direct relation between d_f and n is not yet clear in those systems.¹⁰⁰

Another important factor is viscous coupling and hydrodynamics. In polymeric systems, hydrodynamic interactions lead to a different expression of n as a function of d_f .⁹⁵ Through normal mode analysis of weak colloidal gels it has been shown that hydrodynamic interactions can drive a power-law relaxation response.¹⁰¹ Finally, the viscous coupling of spring networks with solvents have also been shown to affect the value of n .¹⁰² In all of these studies, it has been shown that hydrodynamic interactions result in an increase in n , accelerating the stress relaxation of soft materials.

G. Colloidal gel dynamics

Several studies have also reported unique factors that give rise to the non-Maxwellian relaxation dynamics of colloidal gels. It has been shown in prior work that the density correlations within a single cluster of the gel follows a stretched exponential response.^{103,104} A microrheological conversion¹⁹ of this effect using the generalized Stokes–Einstein approach results in a Kelvin–Voigt like mechanical response, wherein the material acts like a solid at long times (low ω) and a liquid at short times (high ω). In such permanent systems there is another collective relaxation mode at longer times that can also manifest as a result of mutual constraints imposed by steric hindrance; this leads to subdiffusive dynamics, and thus a power-law decrease in $G'(\omega)$ and $G''(\omega)$.¹⁰⁵

Zaccone *et al.*, have shown that relating the cluster size distributions arising from attractive gelation of colloids to the continuous relaxation spectrum $H(\tau)$ can result in different relaxation spectra based on the fractal dimension of the associative colloidal gel (for instance, from gelation that proceeds *via* reaction-limited aggregation or diffusion-limited aggregation). Using this approach, a stretched exponential form of the continuous relaxation spectrum $H(\tau)$ (see additional detail on $H(\tau)$ in Section IVA) is obtained for diffusion-limited aggregation, and a power-law $H(\tau)$ is obtained for reaction-limited aggregation and chemical gelation.¹⁰⁶

H. Non-linearity, non-affinity, and intermittency

A popular approach to understand non-Maxwellian relaxation in soft systems (particularly those that show aging dynamics) is through the soft glassy rheology (SGR) model,^{107,108} in which a power-law relaxation is obtained as a result of activated local yielding processes governed by an exponentially-distributed

energy landscape. A “noise temperature” exponent can be extracted from the exponent of the power-law relaxation x from $G'(\omega) \sim \omega^{x-1}$, which indicates how close the material is to a glass transition. The exponentially-distributed energy landscape is motivated by a trap model used for weakly aging systems.¹⁰⁹ This local yielding process can also be interpreted in terms of shear-transformation zones,^{110–114} which has since become a popular method for interpreting relaxation dynamics of glassy and disordered systems.^{115,116}

Studies have also shown the importance of non-affine deformations on the power-law viscoelasticity of soft matter. This has been explored in particular detail in spring networks, which experience an increase in non-affine deformations near critical connectivity.¹¹⁷ Studies have shown that such non-affine deformations can therefore directly affect the scaling response of the power law exponent governing $G'(\omega) \sim G''(\omega) \sim \omega^n$.^{92,118} Recent works have also shown that such non-affine deformations can also arise in dense suspensions,¹¹⁹ suggesting that the underlying physics governing spring networks and dense colloidal suspensions may be similar. In emulsions, it has also been shown that the power-law response in the loss modulus G'' can arise due to dissipation arising from non-affine motions.¹²⁰

Other studies have also demonstrated the importance of non-linear mechanical driving on the non-Maxwellian rheology of soft materials. For instance, non-linear mechanical stresses can significantly slow down the power-law relaxation observed in semiflexible polymers, with a corresponding decrease in the exponent n in the relation $G(t) \sim t^{-n}$ from $n = 1/2$ as discussed previously in sticky semiflexible polymers, to a value as low as $n = 1/10$.¹²¹ Our recent studies also support the picture of a non-linear mechanical process, where we show that the characteristic relaxation time τ_c of dynamically arrested soft materials such as gels are governed by internal stresses, and that mechanical driving leads to intermittent avalanches which result in a significant broadening in the continuous relaxation spectrum $H(\tau)$.¹²² In this picture, the stretched-exponential-like stress relaxation of dynamically arrested solids are a manifestation of local viscoplasticity arising from the marginal stability of these systems under imposed mechanical deformations.^{123–125}

Studies have also shown that the scaling exponent governing the displacement trajectories of intermittent avalanches in materials that exhibit such marginal behavior (with power-law free energy landscapes) can be quantitatively linked to the scaling exponent of power-law stress relaxation of soft materials. In this framework, the superdiffusive exponent α obtained from the mean-square displacement relation $\Delta r^2(\tau) \sim \tau^\alpha$ can be related to the power-law exponent n obtained from $G(t) \sim t^{-n}$. This requires the knowledge of the power spectrum of stress fluctuations $\langle \Delta \sigma^2(\tau) \rangle \sim \tau^d$; derivations for arrested materials that exhibit spontaneous and random localized motion due to force dipoles (for instance, due to internal stresses^{126,127}) result in $\Delta = 1$,¹²⁸ and a similar value is directly observed in coarsening foams¹²⁹ and cells.¹³⁰ The power-law exponent n can be derived *via* a generalized Stokes–Einstein approach to obtain $n = (a - \Delta)/2$.¹²⁹



III. Introduction to modeling non-Maxwellian relaxations

The various proposed origins of non-Maxwellian relaxation processes in Section II invoke various functional forms of stress relaxation. We now introduce a basic mathematical toolkit to describe these non-exponential relaxation processes.

A. The relaxation spectrum $H(\tau)$

The relaxation time spectrum $H(\tau)$ reveals the underlying relaxation modes governing the stress relaxation response. Each individual relaxation mode is assumed to be exponential, though strictly speaking they can be modified to take other forms, for instance a compressed exponential form.¹²² It is also possible to interconvert $H(\tau)$ into a retardation spectrum $L(\tau)$,¹⁸ and either of these spectra can be used to evaluate $G(t)$ or any other linear viscoelastic functions through the Boltzmann superposition integral (see eqn (5) and (6)).^{131–134} Thus, $H(\tau)$ encodes the distribution of relaxation modes which drive the linear viscoelastic behavior of a system. A Maxwellian system exhibits a single characteristic relaxation time τ_c corresponding to a δ function representation of $H(\tau)$ (Section I), while a non-Maxwellian system will exhibit a broader distribution of $H(\tau)$, thus allowing one to diagnose the statistical origins of non-Maxwellian responses in the rheological responses of the soft material in question.

Deriving the expression for $G(t)$ from a known distribution of $H(\tau)$ is fairly simple, and involves a simple numerical integration of the known functional form of $H(\tau)$. The reverse is not true, however. Obtaining $H(\tau)$ from measurements of $G(t)$ or $G^*(\omega)$ (which are inherently of limited temporal or frequency ranges) is challenging as it requires inverse Laplace transformations. This procedure is ill-posed, meaning that small variations in $G(t)$, including those arising from measurement uncertainties, can cause large variations in $H(\tau)$. This has mandated the use of approximation-based methods to obtain estimates of $H(\tau)$ from experimental data. There are various strategies to perform this operation,^{132,134–136} but the use of Tikhonov regularizations is arguably the most established. This was first demonstrated in problems of viscoelasticity by Honerkamp *et al.*,¹³⁷ but the regularization method is commonly used in other disciplines as well.¹³⁸ The regularization method works by finding the optimal solution for $H(\tau)$ that minimizes the square error of the solution as well as a measure of the roughness of the resulting solution; for the sake of brevity we defer to references for more detailed instruction on the application of the method.^{137–139} Regularization-based methods for obtaining $H(\tau)$ are now readily available for users *via* commercial rheometer software and independent codes.¹³⁹

The relaxation spectrum can be used in the form of discrete modes – in which the relaxation curve is deconstructed into a “line spectrum” of Maxwell modes¹⁴⁰ – or in the form of a continuous curve.^{131,132} Using discrete relaxation modes can be advantageous when the viscoelastic relaxation can be associated directly with well-known underlying relaxation processes, for instance in telechelic metal-coordination-based polymer networks with multiple metal–ligand complexes.¹³ Otherwise, discrete modes can lead to an unnecessarily large number of fitting parameters, as we show in

the next section. A continuous parameterized curve for $H(\tau)$ is a more compact method for studying the underlying relaxation processes arising from a stress relaxation curve, though directly estimating the distribution *a priori* from experimental data can be challenging due to the errors incurred from the inverse Laplace transformation process.^{137,141} Alternatively, fitting the experimental data to common functional forms utilized in rheology (see Sections IV and V) and obtaining their analytical solutions of $H(\tau)$ can sometimes provide a clearer description of the relaxation dynamics that underlie an observed non-Maxwellian viscoelastic response in soft materials.

B. Discrete relaxation spectrum and the generalized Maxwell model

The most straightforward method to model non-Maxwellian relaxation is to assume the presence of discrete Maxwell modes in the relaxation function. A mechanical arrangement in which i Maxwell elements are arranged in parallel gives rise to a Prony series of the Maxwell relaxation equation in eqn (2) such that:

$$G(t) = \sum_{i=1}^N G_i \exp(-t/\tau_i) \quad (12)$$

and

$$H(\tau) = \sum_{i=1}^N G_i \delta(\tau - \tau_i) \quad (13)$$

As discussed previously, this modeling strategy is most useful when there is a known number of relaxation modes. Otherwise, statistical approximations must be made to determine the number of relaxation modes in the system.¹⁴²

A useful heuristic approach is to assume the presence of a relaxation mode per decade of time.¹⁴³ We demonstrate the application of this from the relaxation data of a nanoparticle-crosslinked hydrogel which follows a stretched exponential relaxation behavior (Fig. 4A).¹²² As shown in the data, taking $N = 5$ Maxwell modes results in an excellent fit to the data, but the utility of this method is poor as we now have 10 fitting parameters with no discernible physical meaning. One can employ more sophisticated methods to determine a “parsimonious” relaxation spectrum¹⁴⁴ with the minimal number of modes, for instance by adding a penalty factor for overfitting *via* a Bayesian information criteria.¹⁴⁵ Of course, a purely statistical approach to model selection will be agnostic to the plausibility of the implied physics of the system and therefore must be used with care. There is ongoing research to incorporate such physical insights into statistical models (for instance, increasing the statistical likelihood of models in which the parameters can be restricted to a narrower range of values based on physical insight).¹⁴²

C. Continuous relaxation spectrum

A continuous relaxation spectrum provides a quantitative basis for interpreting the underlying physics of the relaxation phenomenon. Obtaining the continuous relaxation spectrum can be challenging, however, as the regularization process to convert rheological data to $H(\tau)$ as described above can lead to



where τ_c is the characteristic relaxation time (*i.e.*, the most probable value) and σ is the width of the distribution; analytical solutions for different σ values are illustrated in Fig. 3. The log-normal relaxation time spectrum is useful in describing non-Maxwellian relaxation because it represents a direct solution of a Gaussian distribution of activation energies. This function may therefore be useful for describing spatially heterogeneous local activation energy of glasses,^{77,78,165} as well as modeling the polydisperse relaxation of reversible networks.⁷⁶ It may also be useful for modeling systems for which there is a Poisson distribution of relaxation modes, such as multivalent associative gels.⁷⁵

The main challenge with using the log-normal function is the absence of easily implementable analytical solutions for the relaxation modulus or the storage and loss moduli. Therefore, $G(t)$ or $G^*(\omega)$ are usually evaluated by numerical integration of $H(\tau)$ in eqn (15) using eqn (5) and (6);^{145,161,162,166–168} see Fig. 4C and D for demonstration.⁷⁶ We have also included a simple MATLAB protocol for performing these fitting procedures (see Resources).

B. Cole–Cole and fractional Maxwell gel functions

The one-parameter version of the ML function with $z = -(t/\tau_c)^\alpha$ and $a = \alpha$ occurs frequently in the modeling of broadly distributed relaxation phenomena that can be described by fractional differential equations.^{151,154} This ML function provides an analytical solution to the Cole–Cole model, which is a classical model used to describe deviations from the Debye dielectric relaxation model (the dielectric analogue to the Maxwell viscoelastic model), given by $\chi^*(\omega) = 1/(1 + (i\omega\tau_c)^\alpha)$.¹⁶⁹ An exact mathematical analogue to the Cole–Cole model exists in viscoelastic models in the form of the fractional Maxwell gel model, which we introduce in greater detail later in the review.

The analytical description of the Cole–Cole or fractional Maxwell gel relaxation follows:^{150,152}

where the one-parameter Mittag-Leffler function $E_\alpha[-(t/\tau_c)^\alpha]$ interpolates between a stretched exponential relaxation at short times ($t \ll \tau_c$) and a power-law decay in time at long times ($t \gg \tau_c$).¹⁵⁴ It can be shown using a Stieltjes transform (a complex generalization of the Laplace transform) that the relaxation function has a corresponding relaxation spectrum:^{146,151,152}

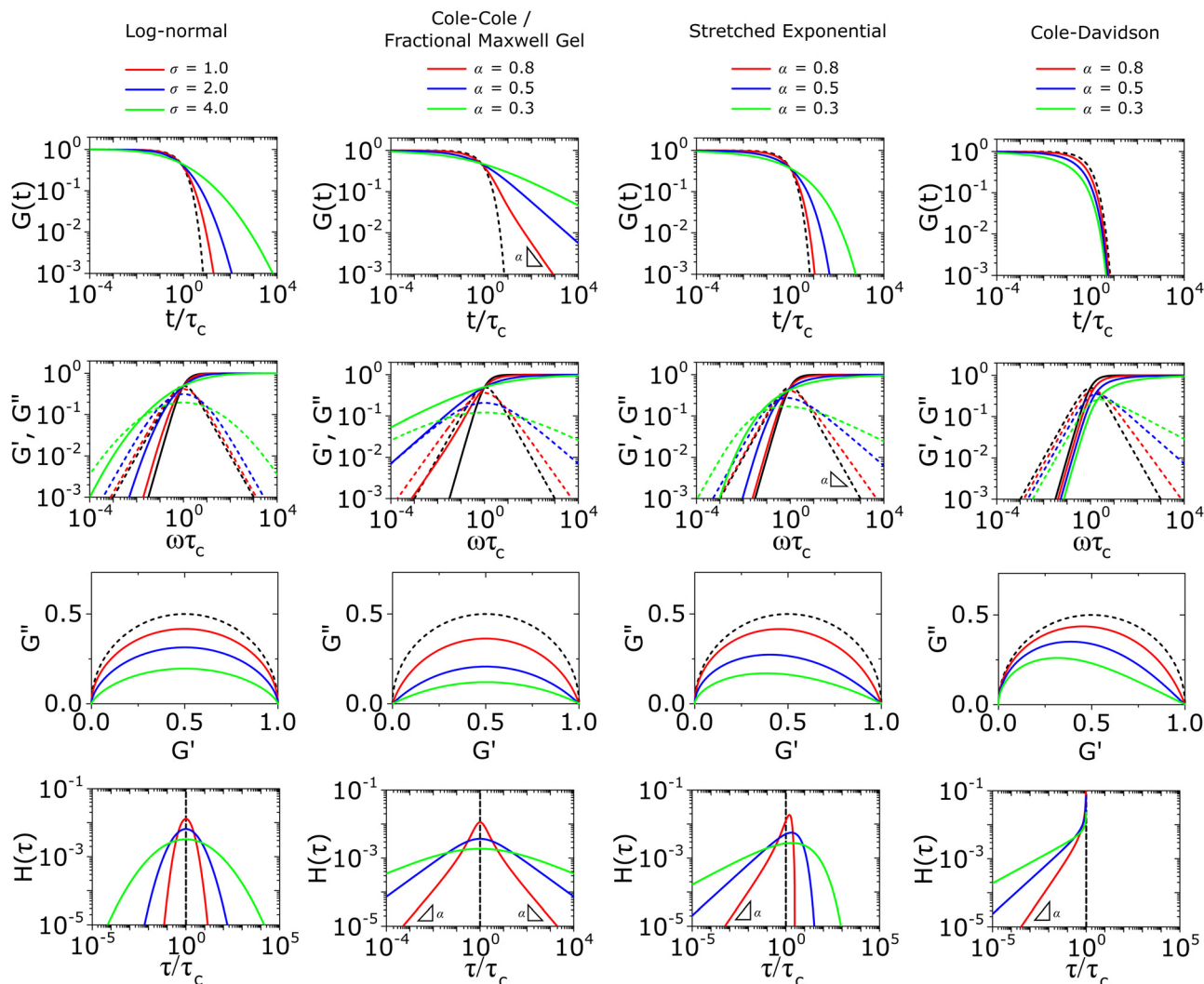
We apply the fundamentals laid out in Section III to highlight common functions used to model non-Maxwellian relaxation processes: the log-normal function, the Cole–Cole or fractional Maxwell gel function, the stretched exponential function, and the Cole–Davidson function. All functions here are valid under normalized conditions such that the initial modulus $G_0 = 1$.

A log-normal distribution of relaxation times has an analytical solution of the form:^{161–164}

which represents a symmetric distribution with power-law tails about the characteristic relaxation time.

The storage and loss moduli can also be derived analytically. The complex shear modulus $G^*(\omega)$ is described by the form:^{150,152,153}

$$G^*(\omega) = G_0 \frac{(\mathrm{i}\omega\tau_c)^\alpha}{1 + (\mathrm{i}\omega\tau_c)^\alpha} \quad (18)$$

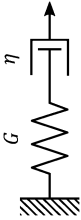
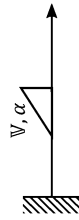
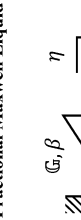




When both the spring and the dashpot are replaced by spring-pots, we obtain the fractional Maxwell model. This represents the most generalized linear viscoelastic model to capture mechanical responses to controlled strain (*e.g.*, $G(t)$, $G'(\omega)$, $G''(\omega)$) as it combines the benefits of the fractional Maxwell gel and the fractional Maxwell liquid models; the fractional Maxwell model can easily be reduced into either of these models by setting one of the fractional exponents to 0 or 1, respectively. The stress relaxation response follows a two-parameter variant of the Mittag-Leffler function in eqn (14), and is useful for modeling viscoelastic materials which exhibit a smooth transition between two power-law regimes. This model is also particularly useful in modeling complex materials such as tissues and food composites.^{190,191} We show a representative example of this in modeling the rheological response of the muscle tissues of Yellowfin tuna, which is more accurately modeled by the fractional Maxwell model than other functions such as the stretched exponential function, the log-normal function, and the generalized Maxwell model (Fig. 6D).

The spring-pot represents the basic building block of fractional mechanical models.^{10,159,160} Individually, it can produce a power-law response in the relaxation modulus (Tables 1, 2 and Fig. 5). The model is thus quite useful in the modeling of critical gels, in which both $G'(\omega)$ and $G''(\omega)$ share a common power-law.⁹⁰ It is noted that critical gels are often also described by a gel strength parameter S ;¹⁸³ this term can be described by $S = \mathbb{V}/\Gamma(1 - \alpha)$ where $\Gamma(x)$ is the complete Gamma function. The spring-pot is a natural choice for modeling critical gels, due to the fractal nature of both the mechanical configuration of the spring-pot as well as the fractal microstructure of critical gels. A spring-pot indeed

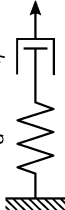
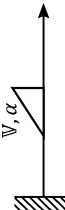
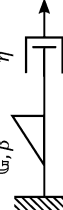
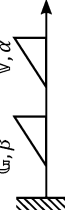
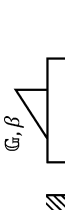
Table 1 Summary of commonly used fractional mechanical models, their constitutive relations, and analytical responses to common rheological deformations (continued)

	Characteristic modulus G_c	Relaxation time τ_c	Constitutive relation	Step strain response $\gamma(t) = \gamma_0 \mathcal{H}(t)$
Maxwell Model				
	$G_c = G$	$\tau_c = \frac{\eta}{G}$	$\sigma(t) + \frac{\eta}{G} \frac{d\sigma(t)}{dt} = \eta \frac{d\gamma(t)}{dt}$	$G(t) = G_c \exp(-t/\tau_c)$
Spring-Pot				
	$G_c \tau_c^\alpha = \mathbb{V}$	$G_c \tau_c^\alpha = \mathbb{V}$	$\sigma(t) = \mathbb{V} \frac{d^{2-\alpha} \gamma(t)}{dt^{2-\alpha}}$	$G(t) = \mathbb{V} \frac{t^{-\alpha}}{\Gamma(1-\alpha)}$
Fractional Maxwell Gel				
	$G_c = \mathbb{V} \tau_c^{-\alpha} \equiv G$	$\tau_c = \left(\frac{\mathbb{V}}{G}\right)^{\frac{1}{1-\alpha}}$	$\sigma(t) + \frac{\mathbb{V}}{G} \frac{d^{2-\alpha} \sigma(t)}{dt^{2-\alpha}} = \mathbb{V} \frac{d^{2-\alpha} \gamma(t)}{dt^{2-\alpha}}$	$G(t) = G_c E_{a,b}(z)$ $a = \alpha$ $b = 1$ $z = -(t/\tau_c)^\alpha$
Fractional Maxwell Liquid				
	$G_c = \eta \tau_c^{-1}$	$\tau_c = \left(\frac{\eta}{G}\right)^{\frac{1}{1-\beta}}$	$\sigma(t) + \frac{\eta}{G} \frac{d^{1-\beta} \sigma(t)}{dt^{1-\beta}} = \eta \frac{d\gamma(t)}{dt}$	$G(t) = G_c (t/\tau_c)^{-\beta} E_{a,b}(z)$ $a = 1 - \beta$ $b = 1 - \beta$ $z = -(t/\tau_c)^{1-\beta}$
Fractional Maxwell Model				
	$G_c = \mathbb{V} \tau_c^{-\alpha} \equiv \left(\frac{G^x}{\mathbb{V}^\beta}\right)^{1/(x-\beta)}$	$\tau_c = \left(\frac{\mathbb{V}}{G}\right)^{\frac{1}{x-\beta}}$	$\sigma(t) + \frac{\mathbb{V}}{G} \frac{d^{x-\beta} \sigma(t)}{dt^{x-\beta}} = \mathbb{V} \frac{d^{2-\alpha} \gamma(t)}{dt^{2-\alpha}}$	$G(t) = G_c (t/\tau_c)^{-\beta} E_{a,b}(z)$ $a = \alpha - \beta$ $b = 1 - \beta$ $z = -(t/\tau_c)^{x-\beta}$
Fractional Kelvin-Voigt Model				
	$G_c = \mathbb{V} \tau_c^{-\alpha} \equiv \left(\frac{G^x}{\mathbb{V}^\beta}\right)^{1/(x-\beta)}$	$\tau_c = \left(\frac{\mathbb{V}}{G}\right)^{\frac{1}{x-\beta}}$	$\sigma(t) = \mathbb{V} \frac{d^{2-\alpha} \gamma(t)}{dt^{2-\alpha}} + G \frac{d^\beta \gamma(t)}{dt^\beta}$	$G(t) = \mathbb{V} \frac{t^{-\alpha}}{\Gamma(1-\alpha)} + G \frac{t^{-\beta}}{\Gamma(1-\beta)}$

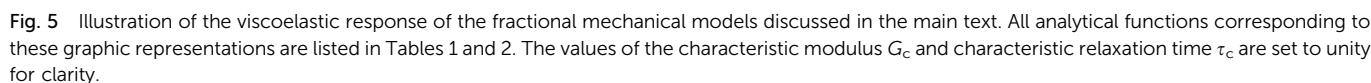
Notations – $G(t)$: stress relaxation modulus; $G'(\omega)$, $G''(\omega)$: storage and loss modulus; $J(t)$: creep compliance; $H(\tau)$: relaxation spectrum. Mathematical functions – delta function $\int_0^{t_0} \delta(x) dx = 1$; the complete Gamma function is denoted $\Gamma(x)$; two parameter Mittag-Leffler function $E_{a,b}(z) = \sum_{n=0}^{\infty} \frac{z^n}{\Gamma(an+b)}$.



Table 2 Summary of commonly used fractional mechanical models, their constitutive relations, and analytical responses to common rheological deformations

	Oscillatory strain response $\gamma(t) = \gamma_0 \sin(\omega t)$	Step stress response $\sigma(t) = \sigma_0$	Continuous relaxation spectrum
Maxwell Model			
	$\frac{G'(\omega)}{G_c} = \frac{(\omega \tau_c)^2}{1 + (\omega \tau_c)^2}$	$J(t) = \frac{t}{\eta} + \frac{1}{G}$	$H(\tau) = G_c \delta(\tau - \tau_0)$
Spring-Pot			
	$G'(\omega) = V \omega^\alpha \cos(\pi \alpha / 2)$	$J(t) = \frac{1}{V} \frac{t^\beta}{\Gamma(1 + \alpha)}$	$H(\tau) = V \frac{\sin \pi \alpha}{\pi \tau^\alpha}$
Fractional Maxwell Gel			
	$\frac{G'(\omega)}{G_c} = \frac{(\omega \tau_c)^{2\alpha} + (\omega \tau_c)^\alpha \cos(\pi \alpha / 2)}{1 + (\omega \tau_c)^{2\alpha} + 2(\omega \tau_c)^\alpha \cos(\pi \alpha / 2)}$	$J(t) = \frac{1}{V} \frac{t^\alpha}{(1 + \alpha)} + \frac{1}{G}$	$H(\tau) = \frac{G_c}{\pi} \frac{(\tau / \tau_c)^\alpha \sin(\pi \alpha)}{(\tau / \tau_c)^{2\alpha} + 2(\tau / \tau_c)^\alpha \cos(\pi \alpha) + 1}$
Fractional Maxwell Liquid			
	$\frac{G'(\omega)}{G_c} = \frac{(\omega \tau_c)^{2-\beta} \cos(\pi \beta / 2)}{1 + (\omega \tau_c)^{2(1-\beta)} + 2(\omega \tau_c)^{1-\beta} \cos(\pi(1-\beta)/2)}$	$J(t) = \frac{t}{\eta} + \frac{1}{G} \frac{t^\beta}{\Gamma(1 + \beta)}$	$H(\tau) = \frac{G_c}{\pi} \frac{(\tau / \tau_c)^{1-\beta} \sin(\pi \beta)}{(\tau / \tau_c)^{\beta-1} + (\tau / \tau_c)^{1-\beta} + 2 \cos(\pi(1-\beta))}$
Fractional Maxwell Model			
	$\frac{G'(\omega)}{G_c} = \frac{(\omega \tau_c)^\alpha \cos(\pi \alpha / 2) + (\omega \tau_c)^{2\alpha-\beta} \cos(\pi \beta / 2)}{1 + (\omega \tau_c)^{2(\alpha-\beta)} + 2(\omega \tau_c)^{\alpha-\beta} \cos(\pi(\alpha-\beta)/2)}$	$J(t) = \frac{1}{V} \frac{t^\alpha}{\Gamma(1 + \alpha)} + \frac{1}{G} \frac{t^\beta}{\Gamma(1 + \beta)}$	$H(\tau) = \frac{G_c}{\pi} \frac{(\tau / \tau_c)^{-\beta} \sin(\pi \alpha) + (\tau / \tau_c)^{-\alpha} \sin(\pi \beta)}{(\tau / \tau_c)^{\beta-\alpha} + (\tau / \tau_c)^{\alpha-\beta} + 2 \cos(\pi(\alpha-\beta))}$
Fractional Kelvin-Voigt Model			
	$G'(\omega) = V \omega^\alpha \cos(\pi \alpha / 2) + G \omega^\beta \cos(\pi \beta / 2)$	$J(t) = \frac{t^\alpha}{V} E_{a,b}(z)$	$H(\tau) = V \frac{\sin \pi \alpha}{\pi \tau^\alpha} + G \frac{\sin \pi \beta}{\pi \tau^\beta}$
	$G''(\omega) = V \omega^\alpha \sin(\pi \alpha / 2) + G \omega^\beta \sin(\pi \beta / 2)$	$a = \alpha - \beta$	
		$b = 1 + \alpha$	
		$z = -\left(-t/\tau_c\right)^{\alpha-\beta}$	

Notations – $G(t)$: stress relaxation modulus; $G'(\omega)$, $G''(\omega)$: storage and loss modulus; $J(t)$: creep compliance; $H(\tau)$: relaxation spectrum. Mathematical functions – delta function $\int_0^{0+} \delta(x) dx = 1$; complete Gamma function $\Gamma(x)$; two-parameter Mittag-Leffler function $E_{a,b}(z) = \sum_{n=0}^{\infty} \frac{z^n}{\Gamma(an + b)}$.

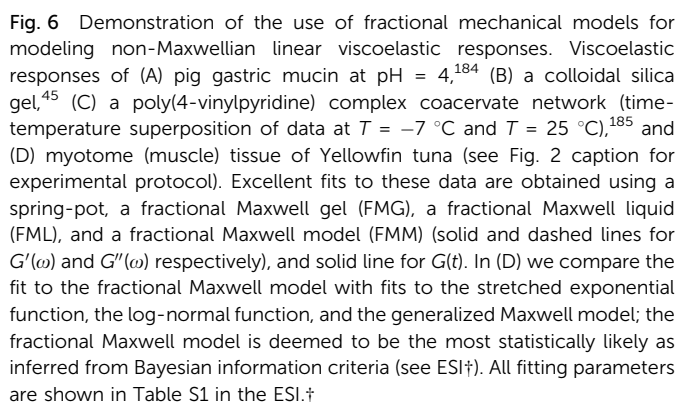


VI. Statistical considerations in modeling rheological data

Using the many models introduced in the tutorial review to fit a given set of rheological data is a process which is inherently governed by statistical methods. When the functional form of a model is known *a priori*, a common method to obtain the “right” parameters for the model that best describes a given set of data is to minimize the weighted residual sum of squares (RSS_w):

$$\text{RSS}_{w_i} = \sum_{i=1}^n \left(\frac{y_i - f(x_i)}{w_i} \right)^2 \quad (26)$$

which computes the sum of the difference between data y and model $f(x)$ which is then scaled by a weighting factor w . The choice of w_i plays an important role in dictating the final parameters of a model, and we illustrate this idea using the Maxwellian viscoelastic data in Fig. 1H. Though the RSS_{w_i} can be used without a weighting factor (*i.e.* $w_i = 1$), since rheological data is often logarithmic, the RSS_{w_i} is conventionally rescaled by the magnitude of the data point such that $w_i = y_i$. However, statistical arguments suggest¹⁹⁴ that the most fundamental weighting factor is $w_i = e_i$, where the absolute error term e_i can be obtained from the absolute uncertainty of measurements arising from a rheometer. Though the latter can be difficult to implement since e_i needs to be measured directly on a rheometer through repeat measurements, Singh *et al.* have recently



When the expected form of the relaxation process is not known *a priori*, one can infer the most likely model for a given set of rheological data using an appropriate statistical information criterion. Though the most straightforward method is to simply minimize the RSS_{w_i} , this method can be prone to overfitting.¹⁴² This overfitting problem can be compensated for by adding a term that penalizes large number of parameters. One such approach is the Bayesian information criterion (BIC):

where $\hat{L}$ is the maximum of the likelihood function, n_p is the number of fitting parameters, and n_i is the number of data points. For data arising from a Gaussian process with a known variance (for instance, e_i), this relation simplifies to:¹⁹⁵

We can obtain the BIC values of the fractional Maxwell model, the generalized Maxwell model (with 5 elements), the stretched exponential function, and the log-normal function used to fit the stress relaxation data of Yellowfin tuna in Fig. 6D (see attached MATLAB demo and Fig. S1 in the ESI[†]). We see that the fractional Maxwell model has the lowest BIC value of the four models, and thus represents the most statistically likely model. A similar operation on the small-amplitude oscillatory shear measurements on the metal-coordinating polymer network of Epstein *et al.* (from Fig. 4C) shows that the log-normal function is the most statistically likely model (Fig. S2, ESI[†]).

These analyses also show that the effect of the penalty term is relatively small compared to the likelihood term; the generalized Maxwell models ($n_p = 10$ for the tuna data and $n_p = 6$ for the metal-coordinating polymer network data) have a lower BIC value than the stretched exponential models ($n_p = 3$) in both cases, despite the stretched exponential exhibiting reasonably good fits to the data. This may be because the e_i values used to assume the absolute error in this analysis – obtained from uncertainties arising from measured variables on the rheometer¹⁹⁴ – provides an under-estimation of the actual errors of the obtained data. More realistic estimations of errors must also consider uncertainties arising from experimental setup (for instance, sample under-filling and overfilling, step strain equilibration time) and from intrinsic heterogeneities in the samples, which would be challenging to quantify. Using larger weighting functions with eqn (27) will lower the magnitude of the likelihood term of the equation and increase the relevance of the penalty term.

We have provided a summary of non-Maxwellian viscoelastic relaxation processes in soft materials, reviewing their diverse origins in different soft materials, and introducing mathematical models and statistical tools to model the observed relaxation responses. The review is aimed at guiding the readers on selecting appropriate mathematical and mechanical models for capturing non-Maxwellian relaxation responses, and drawing meaningful conclusions on the underlying physics of the system based on knowledge of microstructural features of the system (which we have also highlighted in the review). A deep understanding of the principles of soft matter relaxation will have widespread implications in understanding and engineering natural and synthetic soft matter systems, respectively.

We have uploaded a collection of MATLAB codes to demonstrate fitting small-amplitude oscillatory strain and step strain data to continuous relaxation spectra and fractional mechanical models to the File Exchange at <https://www.mathworks.com/matlabcentral/fileexchange/111170-analysis-of-non-maxwellian-viscoelastic-data>. Fitting results using these demonstrations can also be found in the ESI[†] (Fig. S1 and S2).

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